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(54) **Method of preparing a monoclonal antibody, monoclonal antibody, a pharmaceutical composition and a diagnostic reagent**

Verfahren zur Herstellung eines monoklonalen Antikörpers, ein monoklonaler Antikörper, eine pharmazeutische Zusammensetzung und ein diagnostisches Reagenz

Procédé de préparation d'un anticorps monoclonal, un anticorps monoclonal, une composition pharmaceutique et un réactif diagnostique

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Description

[0001] The present invention relates to a method of preparing a monoclonal antibody against a surface antigen, and in particular such a surface antigen which constitutes only an insignificant amount of the total antigen injected in a mammal and a surface antigen which easily loses its in vivo conformation. The problem of an insignificant amount of antigen relative to the total amount of antigen for example sets in when the antigen material used to inject a mammal is derived from another mammalian species.

[0002] The problems associated with the antigens specified above become clear when, for example, it is desired to make monoclonal antibodies against the specificity determining part of a T-cell receptor (TCR). Such (monoclonal) antibodies are known as anti-clonotype antibodies as they recognize the antigen-specific part (or clonotype structure) of a T-cell receptor of one particular T-cell clone. While (murine) monoclonal anti-clonotype antibodies against murine T-cell receptors (TCR) have been reported only occasionally, there are even less (murine) monoclonal anti-clonotype antibodies known against human T-cell receptors. T-cells are hard to culture, making it difficult to obtain and purify a sufficient amount of antigen, i.e. T-cell receptor protein. This in turn makes it difficult to obtain an immune response and also makes screening difficult. In one of the few cases where a monoclonal anti-clonotype antibody is available against a human TCR, these problems did not arise because the antigen was present on Jurkat leukaemia cells (ref. 1). As leukaemia cells can be cultured in unlimited amounts, the problem of a shortage of antigen did not arise. Therefore, until now no method was available to prepare monoclonal antibodies against rare and/or very unstable antigens without having to screen unfavourably large numbers of hybridoma clones.

[0003] The object of the present invention is to provide a method of preparing a monoclonal antibody against a specificity determining part of a cell surface antigen under unfavourable circumstances as described above, said method dramatically reducing the number of hybridoma clones to be screened, rendering the method according to the invention more efficient or even successful where methods according to the state of the art fail.

[0004] To this end the method according to the invention comprises the steps as specified in claim 1.

[0005] For the purpose of keeping the description and the claims readable and intelligible, here the term cell is used not only to indicate mammalian cells but also viruses, and in particular membrane-comprising viruses.

[0006] The term carrier-bound material of cells encompasses intact or whole cells (and viruses), membrane fractions of said intact whole cells (and viruses), or substantially purified surface-antigens or complexes thereof.

[0007] Unless specified otherwise, the term whole cell refers to cells having the surface antigen of interest. The term related cells refers to cells which preferably differ only in that they lack the surface-antigen of interest, or more specifically at least lack the immunological determinant against which it is desired to obtain a monoclonal antibody.

[0008] Surprisingly it proved to be possible to enrich the cell fraction in B-cells specific for the cell surface antigen without the use of cell surface antigen by contacting the isolated B-cells-containing cell fraction with material of cells from the same species as the cell surface antigen-comprising material was derived of but lacking the cell surface antigen. In other words, by contacting the B-cells with irrelevant related cells or cell material and removing bound B-cells an enrichment was achieved.

[0009] Thus it is possible to prepare a monoclonal antibody against a minor cell surface antigen, even if said cell surface antigen suffers from conformational instability.

[0010] European patent EP 0 488 470 and ref. 3, which is based on this European patent, describe a method in which I) a mammal is injected with an antigen, II) a B-cells-containing fraction is isolated, III) B-cells specific for the antigen are selected, IV) the enriched fraction is subjected to clonal expansion and after V) selecting a B-cell clone and immortalisation using a small-scale fusion technique a VI) hybridoma is selected and cloned followed by isolation of a monoclonal antibody comprising fraction. In step III the B-cells are selected by binding them on an antigen coated plastic surface or by rosetting them with antigen coated paramagnetic beads. The non-specific B-cells are removed by washing. Thus, apart from other differences, this method relies on the ample availability and use of antigen - during the selection 2 µg/ml antigen is used to coat plates with antigen -, whereas the present invention solves the problem for a case in which antigen is available in very limited amounts and even highly impure.

[0011] A preferred embodiment is characterized in that the non-human mammal injected with surface antigen is of a different species than the mammalian species from which the surface antigen originates.

[0012] The method according to the present invention is very suitable for preparing monoclonal antibodies when many antigens capable of eliciting an immuneresponse are present.

[0013] This situation exists in particular when the surface antigen has a constant section and a variable section, wherein at least a part of said variable section defines a specificity determining part of said surface antigen.

[0014] According to a preferred embodiment receptor molecule-comprising material is used as the cell surface antigen-comprising material.

[0015] The method according to the invention is particularly suitable for preparing monoclonal antibodies against the specificity determining part of a receptor molecule. The specificity determining part of the receptor molecule is only a minor part of the receptor molecule, which in itself is a minor constituent of all the molecules - and thus antigens - on

the surface of a cell.

[0016] A preferred target according to the invention is a T-cell clone, which T-cell clone is used to prepare the receptor molecule-comprising material.

[0017] A T-cell may be used as a whole cell, or used for the preparation of a membrane fraction thereof. Thus, in the former case the term prepare may, here, relate to the mere isolation of a T-cell or resuspending in a different medium.

[0018] According to preferred embodiments the membrane fraction in step 1 is obtained by mechanical treatment of the whole cells, and the cell surface antigen-comprising material is injected in the mammal in the absence of an adjuvant.

[0019] Both measures help to prevent loss of the in vivo conformation of the surface antigen.

[0020] Furthermore the enriched B-cells-containing cell fraction is preferably further enriched (step 3) by contacting the cell fraction with carrier-bound cell surface antigen comprising material chosen from the group of j) whole cells jj) a membrane fraction obtained from said whole cells and jjj) substantially purified cell surface antigen, and subsequently separating B-cells not bound to said carrier-bound material from B-cells bound to said carrier-bound material, said B-cells bound to said carrier-bound material comprising the further enriched cell fraction.

[0021] This further enrichment can be designated as a positive selection technique, selecting specifically specific B-cells. Thus, though hardly any antigen is available, further enrichment could be achieved.

[0022] According to a preferred embodiment paramagnetic beads are advantageously used as the carrier.

[0023] The use of paramagnetic beads during the enrichment facilitates the separation of wanted and unwanted B-cells.

[0024] An insignificant amount of antigen also presents a problem during screening. According to a preferred embodiment, the selection in at least one of the steps 5 and 6 is conducted using an agglutination assay wherein supernatant of the B-cell clone is contacted with a carrier coated with antibodies capable of binding antibodies of the species of the injected mammal used in step 1 and whole cells bearing the cell surface antigen, and agglutination is detected.

[0025] Simply mixing B-cell culture supernatant, whole cells and a carrier coated with antibodies capable of binding antibodies of the mammalian species used in step 1, allows for a very sensitive and rapid detection of suitable clones while washing is not required.

[0026] Advantageously whole cells related to the whole cells but lacking the surface antigen may be used as a control.

[0027] This allows for the rejection of false-positive clones, and saves time by avoiding superfluous small-scale fusions.

[0028] According to a preferred embodiment of the method according to the invention, the selected B-cell clone in step 5 is mixed with myeloma cells and subjected to mini-electrofusion.

[0029] Mini-electrofusion allows for the efficient fusion of very small numbers (e.g. hundreds) of cells.

[0030] The invention also relates to a pharmaceutical composition comprising a monoclonal antibody prepared according to the invention admixed with a suitable excipient.

[0031] Furthermore the present invention relates to a monoclonal antibody reactive with the clonotypic structure of a T-cell receptor according to claim 11.

[0032] Such monoclonal antibody can be used for diagnostic purposes as well as for the preparation of a pharmaceutical composition.

[0033] More specifically the T-cell receptor is a T-cell receptor associated with an auto-immune disease, and in particular rheumatoid arthritis.

[0034] The monoclonal antibody is reactive with the T-cell receptor of a HC gp-39 reactive T-cell clone, and in particular with the T-cell clone H.243 (ECACC accession No. 96103122).

[0035] Specific examples of suitable monoclonal antibodies are those produced by a hybridoma chosen from the group consisting of TCR 69 (ECACC accession No. 96103118), TCR 70 (ECACC accession No. 96103119), TCR 72 (ECACC accession No. 96103120) and TCR 83 (ECACC accession No. 96103121). These hybridomas have been deposited by NV Organon, PO Box 20, 5340 BH Oss, the Netherlands at CAMR, Porton Down, Salisbury SP4 OJG, UK, in accordance with the Budapest Treaty.

[0036] As indicated above, the invention also relates to pharmaceutical composition comprising a monoclonal antibody as specified in claims 11 to 13 according to the invention admixed with a suitable excipient, suitable for the treatment of rheumatoid arthritis.

[0037] Finally the disclosure relates to the diagnostic use of a monoclonal antibody chosen from the group consisting of a monoclonal antibody prepared using the method according to the invention and a monoclonal antibody according to the invention. Also a diagnostic reagent comprising the antibody as specified in claims 11 to 13 is an embodiment of the invention.

[0038] Now the invention will be explained in further detail with reference to the following example, said example showing the best mode of carrying out the invention applied to the preparation of a murine monoclonal antibody specific for a clonotype of a human T-cell receptor.

LEGENDS TO THE FIGURES

[0039]

Figure 1: Identification of H.243 T-cell surface molecules immunoprecipitated by MAb directed against H.243. SDS/PAGE was performed under non reducing conditions (lane 2, 3 and 4) or under reducing conditions (lane 5, 6 and 7) in a 10% gel. Lane 1: 10 kD ladder marker; lane 2 and 5: control MAb; lane 3 and 6: TCR 83; lane 4 and 7: anti-CD3, OKT3.

Figure 2: Anti-clonotype MAb recognize TCR $\alpha\beta$ in Western blotting.

Immunoprecipitated TCR/CD3 complexes of H.243 were resolved on a 10% SDS-PAGE gel under non reducing conditions and subsequently transferred to PVDF membranes. The membranes were incubated with MAb and finally the bound MAb were detected by using an alkaline phosphatase-conjugated goat anti-mouse Ig second antibody. Lane 1: 10 kD ladder marker, lane 2: TCR 64, lane 3: TCR 66, lane 4: TCR 69, lane 5: TCR 70, lane 6: TCR 72, lane 7: TCR 73, lane 8: TCR 76, lane 9: TCR 78, lane 10: TCR 79, lane 11: TCR 83, lane 12: control MAb, lane 13: medium control.

Figure 3: Immobilized anti-TCR MAb stimulate proliferation of human T-cell clone H.243. Proliferation induced by MAb directed against H.243 (TCR 64 to TCR 83) was compared to control MAb as indicated and to TCR 44 which is an anti-clonotype MAb directed against another TCR. Each value represents the mean counts per 5 min of quadruplicate cultures +/- standard deviation.

Figure 4: Pre-incubation with anti-TCR MAb inhibit or block antigen-driven proliferation of human T-cell clone H.243. a) Inhibition by MAb directed against H.243 (TCR 64 to TCR 83) was compared to control MAb as indicated and to TCR 44 which is an anti-clonotype MAb directed against another TCR. Each value represents the mean counts per 5 min of quadruplicate cultures +/- standard deviation. b) dose response curves of strong inhibiting MAb; TCR 64 (open circles), TCR 70 (closed triangles), TCR 78 (open squares), TCR 83 (closed squares) and control MAb 1 (closed circles). Each value represents the mean counts per 5 min of triplicate cultures.

Figure 5

Anti-clonotype MAb induce anergy in human Th1-clone H.243. Immobilized anti-clonotype MAb or control MAb 1 were incubated overnight with H.243 T cells. After removal of the cells from the anergic stimulus, the cells were given a full stimulus of increasing concentrations of peptide and DRB1*0401 matched APC. Proliferation was assessed by ^3H -thymidine incorporation. Each value represents the mean counts per 5 min of triplicate cultures +/- standard deviation.

Figure 6

Anergic H.243 T cells suppress the response of non-anergic H.243 T cells.

Immobilized TCR 76 or control MAb 1 were incubated overnight with H.243 T cells. Then, both T-cell populations were used in a proliferation assay with increasing concentrations of peptide (or PHA) and APC using 2×10^4 T cells per well. In addition, variable amounts of anergic cells resulting from incubation with TCR 76 were mixed with 2×10^4 non-anergic cells resulting from incubation with Mab 1 and a proliferation assay was performed. A/N: ratio of anergic cells versus non-anergic cells; $1 = 2 \times 10^4$ cells. Proliferation was assessed by ^3H -thymidine incorporation. Each value represents the mean counts per 5 min of triplicate cultures + standard deviation.

EXAMPLES

Materials and methods

Reagents

[0040] Culture medium : DMEM/HAM's F12 (Gibco cat. no. 32500) supplemented with 2500 mg/l sodium bicarbonate, 2.3 mg/l 2-mercaptoethanol, 55 mg/l sodium pyruvate, 1.22 mg/l ethanolamine, 360 mg/l L-glutamine, 4.5×10^{-4} mg/l sodium selenite, 62.5 mg/l sodium penicillin and 62.5 mg/l streptomycin sulphate. In fusion experiments, the medium was further supplemented with 13.61 mg/l hypoxanthine and 3.83 mg/l thymidine. This medium is referred to as DMEM/HAM's F12/HT. Selection of hybridomas was performed in DMEM/HAM's F12/HT supplemented with 1% (v/v) of IL-6 containing supernatant of a human bladder carcinoma cell line T24 (T24CM) and $0.4 \mu\text{M}$ aminopterin.

[0041] Fusion medium: 280 mM inositol, 0.1 mM calcium acetate, 0.5 mM magnesium acetate and 1 mM histidine; specific resistance: $1.11 \cdot 10^4 \Omega \cdot \text{cm}$. Ingredients were dissolved in Milli-Q water and subsequently the conductivity was adjusted to $90 \mu\text{S/cm}$ with Milli-Q water or a solution containing 1 mM calcium acetate and 5 mM magnesium acetate.

Cell cultures

[0042] T-cell clone H.243 was derived from a patient suffering from rheumatoid arthritis. This Th1-like T-cell clone recognizes epitope RSFTLASSETGVG (SEQ ID NO: 1) from Human Cartilage gp-39 (HC gp-39) in the context of DRB1*0401 (belongs to MHC class II). The TCR of this clone is characterized as V α 8 and V β 9 positive. Cells were routinely cultured in DMEM/HAM's F12 supplemented with 10% FCS, 20 U/ml IL-2, 5 U/ml IL-4 and periodically restimulated with antigen and histocompatible Antigen Presenting Cells (APC) or with phytohemagglutinin (PHA) and feeder cells. For proliferation assays Foetal Calf Serum (FCS) was replaced by Normal Human Serum (NHS).

Preparation of T-cell lysate and loading beads with TCR/CD3 complexes

[0043] Ten to fourteen days after restimulation, T-cells were washed with PBS and solubilized by incubating (30 min, 0°C) them at a concentration of 10⁸ cells/ml in extraction buffer according to Oettgen et al. (ref. 2) (10 mM triethanolamine, 150 mM NaCl, 1 mM Na₂EDTA, 1% digitonin, 10 μ g/ml leupeptin, 10 μ g/ml aprotinin, 1 μ g/ml pepstatin, 1 mM Pefabloc® AEBSF and 1,8 mg/ml iodoacetamine pH 7.8). The mild detergent digitonin allows extraction of intact TCR/CD3 complexes. Nuclear and cellular debris was removed by centrifugation at 16,000 g for 15 min. at 4°C and the supernatant was stored in aliquotes at -20°C until use for selection of B-cells or immunoprecipitation.

[0044] For selection of specific B-cells, paramagnetic beads were loaded with TCR/CD3 complexes.

[0045] Briefly, 100 μ l Sheep anti-mouse Ig coupled paramagnetic beads (SAM-beads; Dynabeads® 110.02) were incubated (overnight, 4°C) with 22 μ g anti-CD3, OKT3, in PBS with 0.1% BSA. After washing the beads with PBS/BSA, 1.4 x 10⁸ cell equivalents of T-cell lysate and an equal volume (1.4 ml) PBS/BSA with 1% normal mouse serum were added. The latter was added to prevent binding of B-cells to free SAM binding sites on the beads. The bead suspension was incubated for 2 to 4 hours at 4°C and washed with PBS/BSA before use.

Immunization

[0046] Six-week old female BALB/C mice were immunized at 3-to 7 week intervals with either 5 x 10⁶ whole T-cells (invention) or paramagnetic beads loaded with TCR/CD3 complexes of 5 x 10⁷ cell equivalents (control experiment). Different routes of administration were used as indicated in Table I. In case adjuvant was used (control experiment), the first injection was performed with complete Freund's adjuvant, subsequent injections with incomplete Freund's adjuvant and the final boost without adjuvant.

Generation of anti-clonotype MAb

[0047] Female BALB/C mice, 6 weeks of age, were injected intraperitoneally 4 times at 3- to 7-week intervals, each time with 5 x 10⁶ resting T-cells in PBS. Five days after the final injection, mice were sacrificed and erythrocyte and monocyte-depleted spleen cell populations were prepared as described previously (refs. 3, 4). This results in a cell fraction enriched in B-cells, but does not result in the enrichment of B-cells specific for said cell surface antigen with respect to other non-specific B-cells according to step 2 of the present invention.

[0048] To preclear these spleen cell populations for B-cells reactive to human CD3 or the constant region of TCR $\alpha\beta$, approximately 3 x 10⁷ cells were incubated twice with 20 μ l SAM beads loaded with TCR/CD3 complexes of an irrelevant T-cell clone (V α 3.V β 14 positive T-cell clone, SCRO.08A, but any other irrelevant T-cell will do). Subsequently, B-cells specific for the variable region of the TCR were selected by incubating the resulting cell suspension with SAM beads loaded with TCR/CD3 complexes of the T-cell clone of interest (H.243). Each incubation was performed for 90 min. at room temperature in DMEM/HAM's F12 supplemented with 10% Calf Serum (Hyclone®) and 0.2% normal murine serum. During these incubations, the suspension was carefully homogenized every 5 min. After the final incubation, the beads were carefully washed five times with DMEM/HAM's F12, 10% Calf Serum and resuspended in the same medium.

[0049] Monoclonal antibody producing hybridomas were generated from these selected B-cells by clonal expansion and mini-electrofusion as described previously (3). Briefly, selected B-cells were mixed with T-cell supernatant (TSN) and 50,000 irradiated (2500 rad) EL-4 B5-cells at a final volume of 200 μ l DMEM/HAM's F12 supplemented with 10% FCS in 96-well flat bottomed tissue culture plates. At day 8, supernatants were tested in a one-step T-cell agglutination assay as described below using either the T-cell clone of interest or an irrelevant T-cell clone. B-cell cultures producing discriminating MAb were submitted to a mini-electrofusion procedure. The contents of these B-cell cultures were mixed with 10⁶ NS-1 myeloma cells and rendered serum-free by washing with DMEM/HAM's F12/HT. Then, the cells were treated with pronase solution for 3 min. and subsequently washed with fusion medium. Electrofusion was performed in a 50 μ l fusion chamber by an alternating electric field of 30 s, 2 MHz, 400 V/cm followed by a square, high field pulse of 10 μ s, 3 kV/cm and again an alternating electric field of 30 s, 2 MHz, 400 V/cm. Finally, the contents of the fusion chamber was transferred to 20 ml selection medium and plated into a 96-well microtiter plate. At day 8 after fusion, the

cultures were examined for hybridoma growth and screened in the one-step T-cell agglutination assay again.

T-cell agglutination assay

[0050] Screening of hybridoma cultures for anti-T-cell antibodies and determination of cross-reactivity of MAb with other T-cells was performed in a one-step agglutination assay. Using half-area microtiter plates 50 μ l of hybridoma supernatant was mixed with 20 μ l bead/cell suspension containing 2×10^5 paramagnetic beads with covalently bound Sheep anti-Mouse Ig and 5×10^4 T-cells of interest in DMEM/HAM's F12. After 2 to 3 hours incubation at 37°C, agglutination was examined under a microscope by looking for aggregates of cells and beads.

Immunoprecipitation and Western blotting

[0051] Ten to fourteen days after restimulation, T-cells were washed and cell surface proteins were labelled with biotin. Briefly, T-cells were incubated at a concentration of 5×10^6 cells/ml with 0,5 mg/ml ImmunoPure® sulfo-NHS-biotin in PBS for 30 min. at room temperature. Subsequently, the cells were washed again and solubilized as described above. Before use in an immunoprecipitation experiment, the cell lysate was precleared once by incubating with SAM-paramagnetic beads. Immunoprecipitation was performed by incubating 10^7 cell equivalents of precleared lysate for 3 - 4 h with 15 μ l SAM-paramagnetic beads preloaded with MAb (1 ml hybridoma supernatant). Then, immunoprecipitates were washed three times with PBS, boiled in sample buffer and submitted to SDS-PAGE on a 10% gel either under reducing conditions or non-reducing conditions, using the method of Laemmli (ref. 5) and the Mini-protein II system (Biorad). Immunoblotting was performed according to Towbin et al. (ref. 6) using PVDF membranes. Non-specific binding sites were blocked by incubating the membranes with 5% skim milk in PBS supplemented with 0.5% Tween 20. After washing, the blots were probed with streptavidine-alkaline phosphatase in PBS, 0.5% Tween 20, 1% BSA, 1% normal goat serum for 1 h at room temperature. Finally, the blots were developed with BCIP and NBT as a chromogenic substrate. For Western blot studies, paramagnetic SAM-beads were loaded with TCR/CD3 complexes as described above using OKT3 as the immunoprecipitating antibody and normal T-cell lysate. After SDS-PAGE on a 10% PAA-gel under reducing or non-reducing conditions, the blots were incubated with 2.5 ml hybridoma supernatant. Bound antibodies were detected using alkaline phosphatase-conjugated goat anti-mouse Ig and BCIP/NBT as described above.

Antibody mediated T-cell proliferation

[0052] Flat-bottomed microtiter wells were coated (overnight, room temperature) with 100 μ l Sheep anti-mouse Ig at 40 μ g/ml in PBS. The wells were washed once and then incubated with 100 μ l MAb at a concentration of 2 μ g/ml in PBS for 2 h. Excess of free MAb was removed by washing and 2.5×10^4 resting T-cells were added in 200 μ l DMEM/HAM's F12 supplemented with 10% NHS. After two days incubation at 37°C, the cells were pulsed with 0.5 μ Ci [3 H]-thymidine and incubated for another 16 to 18 h. Finally, the cells were harvested onto glass fibre filters and [3 H]-thymidine incorporation was measured by beta counting on a Matrix 96™ (Packard). Each variable was tested in quadruplicates.

Antibody mediated inhibition of antigen-induced proliferation

[0053] Flat-bottomed microtiter wells are inoculated with 20 μ l hybridoma supernatant, 2×10^4 resting T-cells and 1×10^5 histocompatible MNC in 150 μ l DMEM/HAM's F12 supplemented with 10% NHS. After 3 h incubation at 37°C, 50 μ l of peptide RSFTLASSETGVG (16 μ g/ml) was added to the cultures. The cultures were incubated for another two days at 37°C and [3 H]-thymidine incorporation was determined as described above. Each variable was tested in quadruplicate.

Induction of anergy using clonotype-specific MAb

[0054] Twenty-four well culture plates were coated (overnight, room temperature) with 1 ml Sheep anti-mouse Ig at 40 μ g/ml in PBS. The wells were washed once and then incubated with 1 ml MAb at a concentration of 2 μ g/ml in PBS for 2 h. Excess of free MAb was removed by washing and 2×10^6 washed, resting T cells were added in 2 ml DMEM/HAM's F12, 10% NHS. Where indicated cyclosporin A (Sandoz, Basel Switzerland), rIL-2, cycloheximide (Sigma, St Louis, MO, USA) or HLA-DRB1*0401 matched APC (3000 rad irradiated) were added at the time of culture initiation. After overnight incubation, the plates were chilled on ice and the T cells were resuspended by pipetting. The cells were washed once with complete culture medium and used for a T-cell proliferation assay, cytokine analysis and FACS analysis as described below.

The antigen-specific proliferative response of T cells was assessed in flat-bottomed microwell cultures containing 2×10^4 T cells, 1×10^5 HLA-DRB1*0401 matched, irradiated (3000 rad) PBMC and variable antigen concentrations in 200

μl DMEM/HAM's F12, 10 % NHS. After two days incubation at 37 °C, the cells were pulsed with 0.5 μCi [3H]-thymidine and incubated for another 16-18 hours. Finally, the cells were harvested onto glass fiber filters and [3H]-thymidine incorporation was measured by gasscintillation on a Matrix 96 (Packard, Meriden, CT, USA). Each variable was tested in triplicate.

Results

Immunisations

[0055] The immunisations using adjuvant (control experiment) did not yield any sera containing antibodies capable of discriminating between H.243 and irrelevant T-cells (Table 1; Immun. group II, III, and IV). The titer of the sera of group V were low (1:100 in the agglutination assay), whereas those of group I and II were much higher (1:1200).

Generation of anti-clonotype MAb

[0056] Using spleen cells from group I, incubation with TCR/CD3 complexes from an irrelevant T-cell clone, here H. 258, loaded paramagnetic beads resulted in many rosetted cells, indicating a successful removal of many CD3- and TCRαβ-reactive B-cells. When non-rosetted cells were submitted to a second preclear with irrelevant TCR/CD3-complexes from the same T-cell clone, hardly any rosetted cells were visible.

[0057] Microscopic examination after incubation with paramagnetic beads with the T-cell clone of interest, did not result in visible rosette formation. Given the expected frequency of specific B-cells, this is not surprising. After limiting dilution and clonal expansion 36 B-cell supernatants were found to agglutinate H.243 and not the irrelevant T-cell clone H.258 (Table II). Some B-cell supernatants were able to agglutinate both clones, indicating that they had escaped the preclear procedures.

[0058] Further, it can be deduced from Table II that the major part of the specific B-cell cultures are not clonal as also B-cells had grown out that do not react with any of the T-cells. However, this was not a problem because hybridomas from these non-specific B-cells could be selected out after mini-electrofusion. Eighteen of the specific B-cell cultures were submitted to a mini-electrofusion. Some fusions did not result in T-cell specific hybridomas and some others could not be cloned to stable cell lines. Ultimately stable hybridomas were obtained in 11 out of 18 mini-electrofusions. These 11 MAb were further characterized. The isotypes of these MAb are shown in Table III.

Specificity of MAb

[0059] To further investigate the specificity of the MAb, immunoprecipitation and T-cell agglutination tests as well as Western blots were performed.

[0060] SDS/PAGE under non-reducing and denaturing conditions showed that all MAb were able to immunoprecipitate a major band of approximately 85 kD and a minor band of 21 kD from a digitonin lysate of T-cell clone H.243. This is consistent with the molecular weight of the TCRαβ-complex and of CD3 chains ε/δ, respectively. Under reducing conditions the 85kD band dissociates into two bands of 40 and 50 kD which is consistent with molecular weight of the TCR α- and β-chains, respectively. The immunoblot of one of the anti-H.243 specific MAb is shown in Figure 1, together with OKT3 as a positive control and an irrelevant MAb as a negative control.

[0061] Table III shows cross-reactivity of the MAb with six Vβ9 positive T-cell clones, two Vα8 positive T-cell clones and eight other T-cell clones with different Vα and Vβ's. Antibody TCR 74 should be regarded as Vβ9 specific as it agglutinates all the Vβ9 positive T-cells and not the others. The other TCR antibodies only react with H.243 so it is very likely that these antibodies are directed against the antigen-binding part of H.243 as reaction with Vα8, Vβ9, constant region of TCRαβ, CD3 and other common T-cell surface molecules is excluded. Figure 2 shows that antibodies from the hybridoma clones TCR 64, TCR 66, TCR 69, TCR 70, TCR 73, and TCR 76 MAb are able to stain the TCR on a non-reduced Western blot (85 kD band). The non-specific heavy bands at the top of the figure are descended from the immunoprecipitating antibodies SAM and OKT3. No staining with the anticlonotype MAb was obtained in a Western blot ran under reducing conditions (results not shown), suggesting that the MAb recognize a conformation epitope formed by the intact TCRαβ complex.

MAb induced T-cell proliferation

[0062] To investigate whether the MAb are able to induce proliferation of H.243 T-cells, MAb were immobilized on flat bottomed microtiter plates and incubated with T-cells. Though there are great differences between the MAb, Figure 3 shows that all of them can induce proliferation of H.243 cells. Proliferation similar to anti-CD3 (OKT3) was obtained with TCR 64, TCR 66, TCR 70, TCR 76 and TCR 79. No proliferation was obtained with TCR 44, which is an anti-clonotype

MAB for another TCR.

MAB mediated inhibition of antigen-induced proliferation of T-cells

[0063] Further it was investigated whether the panel of MAB could inhibit antigen-driven proliferation of H.243 T-cells. In this experiment, MAB were pre-incubated with T-cells and APC and three hours later, pulsed with different concentrations of peptide. Figure 4 shows that TCR 64, TCR 70, TCR 76, TCR 78 and TCR 83 strongly inhibit antigen-driven proliferation. Up to 98% inhibition was observed with TCR 83. Dose response curves show that maximal inhibition could be obtained with as little as 100 ng/ml TCR 64, TCR 70 and TCR 83. TCR 78 was approximately tenfold less potent. The inhibition patterns were not dependent of whether suboptimal or saturating concentrations of peptide were added (results not shown). An anti-clonotype MAB directed against an other TCR (TCR 44) did not show any inhibition.

Anti-clonotype MAB induce unresponsiveness of human T-cell clone H.243 to subsequent stimulation with antigen and APC

[0064] T-cell receptor occupancy in the absence of costimulation is known to induce anergy. Previously we had found that immobilized anti-clonotype MAB to H.243 can functionally trigger the TCR of this clone. Therefore, it was interesting to investigate whether the same antibodies could induce anergy. For this purpose, 10 different anti-clonotype MAB were immobilized on 24-well plates and incubated overnight with H.243 T cells to give the anergic stimulus. Then, T cells were removed from the plates and it was tested whether they could respond to increasing concentrations of HC gp-39²⁶²⁻²¹⁷ (SEQ ID NO: 3) presented by irradiated, DRB1*0401-matched APC. Figure 5 shows that this response was completely abrogated by 8 out of 10 MAB whereas H.243 cells incubated with control MAB 1 still responded well to peptide presented by APC. The response of T cells incubated with TCR 69 and TCR 83 was significantly reduced but not completely abrogated at higher peptide concentrations.

Anergic cells suppress the response of non-anergic cells

[0065] It was also investigated whether anergic H.243 cells were able to suppress the response of non-anergic H.243 cells. Anergy was induced with TCR 76 and non-anergic cells were obtained from incubation with control MAB 1. Subsequently, proliferation assays were performed with peptide HC gp-39²⁶¹⁻²⁷⁵ (SEQ ID NO: 2) using different T-cell populations viz. 2×10^4 anergic T cells per well, 2×10^4 non-anergic T cells per well or mixtures of decreasing concentrations of anergic T cells (4×10^4 , 2×10^4 , 1×10^4 and 0.5×10^4) and 2×10^4 non-anergic T cells per well. The response of non-anergic cells was significantly reduced in a dose related fashion by the addition of anergic cells (Figure 6). Approximately 90% reduction of proliferation was obtained at an antigen concentration of $1 \mu\text{g/ml}$ and a ratio anergic cells versus non-anergic cells of 2:1. At higher antigen concentrations less reduction of proliferation was observed (72% at $5 \mu\text{g/ml}$ and 39% at $25 \mu\text{g/ml}$).

Thus this example shows that it is possible to generate anti-clonotype monoclonal antibodies against human T-cell receptors. This experiment resulted in ten hybridomas, four of which were deposited with the ECACC (Salisbury, UK), as indicated in Table III. The strategy did not require large numbers of T-cells or ample amounts of purified TCR. The method according to the invention is an attractive alternative for tedious recombinant procedures in which soluble TCR of TCR expressed on syngeneic mouse cells are generated to be used in immunization procedures. Thus also possible problems with conformational stability are avoided. As the monoclonal antibodies showed different responses in a test for antibody induced T-cell proliferation and a test for antibody mediated inhibition of antigen-driven T-cell proliferation (up to 98% inhibition was observed), this suggests that different epitopes are recognized on the same clonotypic structure.

[0066] If cross-linked all anti-clonotype monoclonal antibodies were able to induce proliferation of H.243 T-cells. It is demonstrated that small amounts of immobilized anti-clonotype MAB can induce anergy in the autoreactive clone. Following the anergic stimulus, T cells failed to proliferate upon restimulation as a result of a lack of IL-2 gene transcription. In addition, a diminished IFN γ production was found.

FACS analysis of the anergic cells indicated that anergy was not a result of TCR downmodulation or the absence of free TCR. In coculture experiments, anergic T cells were found to suppress the response of reactive cells from the same clone. This bystander suppression led to 90% inhibition of proliferation. The data demonstrate the anergizing potential of clonotype-specific antibodies in vitro. Such MAB can be used for the induction and maintenance of antigen-specific T-cell tolerance in rheumatoid arthritis.

[0067] In sera of mice immunized by injecting rather pure TCR/CD3 immune complexes coupled to paramagnetic beads (Table I; immunization group IV). The sera of the mice tested more positive for the specific T-cell which was used for immunization than an irrelevant T-cell. However, it was not possible to obtain hybridomas producing anti-clonotype monoclonal antibodies. Even though in sera of mice immunized with small amounts of whole T-cells no specificity for the T-cell of interest versus an irrelevant T-cell could be demonstrated, the method according to the invention yielded

hybridomas capable of producing anti-clonotype monoclonal antibodies.

[0068] It goes without saying that antigen binding fragments can be made of the monoclonal antibodies according to the invention, for example using papain. Also the monoclonal antibodies or antigen binding fragments thereof can be labelled to facilitate their detection and scope of the present invention encompasses all these forms.

5 **[0069]** It also goes without saying that the irrelevant cells used to remove non-specific B-cells, are preferably as much alike as the cell surface antigen-carrying cells.

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Table I: Immunization schedules and characterization of sera

Immun group	antigen	adjuvant	route	agglutination/FACS T cell 1* (immun.) T cell 2* (irrelevant)	proliferation induction	inhibition Ag-driven proliferation [#]	West blot
I	T cells	none	ip (5x)	+++	-	++	-
II	T cells; TCR/CD3	yes	ip;im;im;ip	+++	-	++	-
III	TCR/CD3	yes	ip;im;im;im; ip	-	+/-	-	-
IV	TCR/CD3	yes	sc;sc;sc;ip	-	+	-	-
V	TCR/CD3	none	is (5x)	+	+	-	-
	Normal Mouse Serum			-	-	-	-

Groups of two BALB/C mice were immunized with either whole T cells or TCR/CD3 complexes bound to paramagnetic beads. Immunizations were performed in the absence or presence of adjuvant as indicated. Different routes of immunization were followed namely: ip = intraperitoneally; im = intramuscular; sc = subcutaneously; is = intrasplenic.

Sera from the mice were characterized for binding in a T-cell agglutination assay, FACS-staining and Western blot. Functional characterization was performed in an antibody-mediated T-cell proliferation assay and in a test for antibody-mediated inhibition of antigen-driven T-cell proliferation.

* T cell 1= T-cell clone used for immunization; T cell 2 = irrelevant T-cell clone.

[#] + means inhibition of T-cell proliferation.

Table II: Outgrowth of anti-T cell specific B cells in EL-4 B-cell cultures

Number of wells inoculated	Plating density	Number of wells with B-cell outgrowth	Agglutination with H.243	Agglutination with H.258
384	x	384	68	40
192	1/6 x	171	13	7
192	1/36 x	81	3	1

28 x 10⁶ lymphocytes were submitted to an immunobead selection procedure for the selection of anti-clonotype specific B cells. The selected B cells were expanded at different plating densities in an EL-4 B-cell culture system and at day 8, supernatants were tested for agglutinating antibodies with the T cells of interest (H.243) and irrelevant T cells (H.258)
x: undetermined number of selected cells

Table III: Agglutination of different kinds of T cells with anti-TCR mAb

mAb	isotype	T-cell agglutination with			
		H.243 ¹⁾	6 Vβ9 positive T cells	2 Vα8 positive T cells	8 other T cells
TCR 64	IgG2a	+	-	-	-
TCR 66	IgG1	+	-	-	-
TCR 67	n.d.	+	-	-	-
TCR 69	IgG1	+	-	-	-
TCR 70	IgG2a	+	-	-	-
TCR 72	IgM	+	-	-	-
TCR 73	IgG1	+	-	-	-
TCR 74	n.d.	+	+	-	-
TCR 76	IgG2a	+	-	-	-
TCR 78	IgM	+	-	-	-
TCR 79	IgG1	+	-	-	-
TCR 83	IgM	+	-	-	-
anti-Vβ9	IgG2a	+	+	-	-
anti-αβ	IgG2b	+	+	+	+
control mAb	IgG1	-	-	-	-

1) H.243: Vα8 en Vβ9 positive

LITERATURE

[0070]

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2. Oetgen, H.C. et al. (1986) A T3-like protein complex associated with the antigen receptor on murine T cells. *Nature* **320**: p. 272-275.
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hybridomas. J. Immunol. Methods **152**, p. 69-77.

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Abbreviations

[0071]

PBS Phosphate Buffered Saline
BCIP 5-bromo-4-chloro-3-indoylphosphate p-toluidine salt
NBT p-nitro blue tetrazolium chloride
BSA Bovine Serum Albumin
FCS Foetal Calf Serum
SAM Sheep anti Mouse
TSN T-cell supernatant
MAb monoclonal antibody (antibodies)

Manufacturers

[0072]

Biorad, Richmond, CA, U.S.A.
Gibco, Paisley, Scotland
Hyclone, Logan, U.S.A.
Packard, Meriden, CT, U.S.A.

SEQUENCE LISTING

[0073]

(1) GENERAL INFORMATION:

(i) APPLICANT:

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(ii) TITLE OF INVENTION: Method of preparing a monoclonal antibody, monoclonal antibody, a pharmaceutical composition and a diagnostic reagent

(iii) NUMBER OF SEQUENCES: 3

(iv) COMPUTER READABLE FORM:

(A) MEDIUM TYPE: Floppy disk
(B) COMPUTER: IBM PC compatible
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
(D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)

(2) INFORMATION FOR SEQ ID NO: 1:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 13 amino acids
 (B) TYPE: amino acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(v) FRAGMENT TYPE: internal

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

Arg Ser Phe Thr Leu Ala Ser Ser Glu Thr Gly Val Gly

1

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(2) INFORMATION FOR SEQ ID NO: 2:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 15 amino acids
 (B) TYPE: amino acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(v) FRAGMENT TYPE: internal

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Phe Gly Arg Ser Phe Thr Leu Ala Ser Ser Glu Thr Gly Val Gly

1

5

10

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(2) INFORMATION FOR SEQ ID NO: 3:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 16 amino acids
 (B) TYPE: amino acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(v) FRAGMENT TYPE: internal

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

Gly Arg Ser Phe Thr Leu Ala Ser Ser Glu Thr Gly Val Gly Ala Pro

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Claims

1. Method of preparing a monoclonal antibody against a specificity determining part of cell surface anogen having a constant section and a variable section, wherein at least a part of said variable section defines a specificity determining part of said surface antigen, comprising the steps of

1) injecting a non-human mammal with cell surface antigen-comprising material, said material being chosen from the group consisting of i) whole cells and ii) a membrane fraction obtained by treating whole cells;
 2) isolating a B-cells-containing cell fraction from the spleen of said mammal;
 3) enriching said cell fraction obtained in step 2 in B-cells specific for said cell surface antigen by contacting the cell fraction with carrier-bound material of cells related to said whole cells, said carrier-bound related material of cells lacking said cell surface antigen, and separating B-cells bound to carrier-bound material of related cells from the enriched unbound B-cells-containing cell fraction to be used in the next step;
 4) contacting the cell fraction with carrier-bound cell surface antigen comprising material chosen from the group of j) whole cells and jj) a membrane fraction obtained from said whole cells and subsequently separating B-cells not bound to said cancer-bound material from B-cells bound to said carrier-bound material,
 5) subjecting the B-cells bound to said carrier-bound material obtained in the previous step to limiting dilution followed by clonal expansion;
 6) selecting a B-cell clone and immortalising said selected B-cell clone using a small-scale fusion technique: and
 7) selecting and cloning a hybridoma capable of producing antibodies which specifically bind said cell surface antigen, followed by isolating a monoclonal antibody-comprising fraction from supernatant of said hybridoma.

2. Method according to claim 1, **characterized in that** the mammal injected with surface antigen is of a different species than the mammalian species from which the surface antigen originates.

3. Method according to claim 2, **characterized in that** receptor molecule-comprising material is used as the surface antigen-comprising material.

4. Method according to claim 3, **characterized in that** a T-cell done is used to prepare the receptor molecule-comprising material.

5. Method according to any of the preceding claims, **characterized in that** the membrane fraction in step 1 is obtained by mechanical treatment of the whole cells.

6. Method according to any of the preceding claims, **characterized** the cell surface antigen-comprising material is injected in the mammal in the absence of an adjuvant.

7. Method according to any of the preceding claims, **characterized in that** paramagnetic beads are used as the carrier.

8. Method according to any of the preceding claims, **characterized in that** the selection in at least one of the steps 6 and 7 is conducted using an agglutination assay wherein supernatant of the B-cell clone is contacted with a carrier coated with antibodies capable of binding antibodies of the species of the injected mammal used in step 1 and whole cells bearing the cell surface antigen, and agglutination is detected.

9. Method according to claim 8, **characterized in that** whole cells related to said whole cells but lacking said cell surface antigen are used as a control.

10. Method according to any of the preceding claims, **characterized in that** the selected B-cell clone in step 6 is mixed with myeloma cells and subjected to mini-electrofusion.

11. Monoclonal antibody reactive with the clonotypic structure of a T-cell receptor, **characterized in that** said monoclonal antibody is reactive with the T-cell receptor of a HC gp-39 reactive T-cell clone.

12. Monoclonal antibody according to claim 11, **characterized in that** the T-cell clone is ECACC accession No. 96103122.

13. Monoclonal antibody according to claim 11, **characterized in that** it is produced by a hybridoma chosen from the group consisting of ECACC accession No. 96103118, ECACC accession No. 96103119, ECACC accession No. 96103120 and ECACC accession No. 96103121.

14. Pharmaceutical composition comprising a monoclonal antibody according to any of the claims 11 to 13 admixed with a suitable excipient, suitable for the treatment of rheumatoid arthritis.

15. Diagnostic reagent comprising a monoclonal antibody according to any of the claims 11 to 13.

Patentansprüche

1. Verfahren zur Herstellung eines monoklonalen Antikörpers gegen einen die Spezifität bestimmenden Teil eines Zelloberflächenantigens mit einem konstanten Abschnitt und einem variablen Abschnitt, wobei mindestens ein Teil des besagten variablen Abschnitts einen die Spezifität bestimmenden Teil des besagten Oberflächenantigens bildet, umfassend die Schritte:

1) Injizieren von Zelloberflächenantigen umfassendem Material in ein nicht-menschliches Säugetier, wobei das besagte Material gewählt worden ist aus der Gruppe, bestehend aus i) ganzen Zellen und ii) einer Membranfraktion, die durch Behandlung von ganzen Zellen erhalten worden ist;

2) Isolieren einer B-Zellen enthaltenden Zellfraktion aus der Milz des besagten Säugetiers;

3) Anreichern der besagten Zellfraktion, die in Schritt 2 erhalten worden ist, in B-Zellen, die für das besagte Zelloberflächenantigen spezifisch ist, durch Kontaktieren der Zellfraktion mit trägergebundenem Material von Zellen, die sich auf die besagten ganzen Zellen beziehen, wobei das besagte trägergebundene bezogene Material von Zellen an dem besagten Zelloberflächenantigen mangelt, und Trennen von B-Zellen, die mit trägergebundenem Material von in Beziehung stehenden Zellen gebunden sind, von der angereicherte ungebundene B-Zellen enthaltenden Zellfraktion, die im nächsten Schritt eingesetzt werden soll;

4) Kontaktieren der Zellfraktion mit dem trägergebundenen Zelloberflächenantigen enthaltenden Material, welches aus der Gruppe j) ganzer Zellen und jj) einer Membranfraktion ausgewählt worden ist, die aus den ganzen Zellen erhalten worden ist, und nachfolgend Trennen der B-Zellen, die nicht an das trägergebundene Material von B-Zellen gebunden sind, von B-Zellen, die an das besagte trägergebundene Material gebunden sind;

5) Unterwerfen der B-Zellen, die mit dem besagten trägergebundenen Material gebunden sind, welches im vorherigen Schritt erhalten worden ist, um die Auflösung, die einer klonalen Expansion folgt, zu begrenzen;

6) Auswählen eines B-Zellen Klon und Immortalisieren des besagten ausgewählten B-Zellen Klon unter Einsatz einer niedrig skalierten Fusionstechnik; und

7) Auswählen und Klonen eines Hybridoms, das fähig ist, Antikörper herzustellen, die in spezifischer Weise das besagte Zelloberflächenantigen anbindet, gefolgt vom Isolieren einer monoklonalen Antikörper enthaltenden Fraktion aus dem Supernatanten des besagten Hybridoms.

2. Verfahren nach Anspruch 1, **dadurch gekennzeichnet, dass** das Säugetier, welchem das Oberflächenantigen injiziert worden ist, von einer anderen Art ist als die Säugetierart, von der das Oberflächenantigen abstammt.

3. Verfahren nach Anspruch 2, **dadurch gekennzeichnet, dass** Rezeptormolekül umfassendes Material als das Oberflächenantigen umfassende Material eingesetzt wird.

4. Verfahren nach Anspruch 3, **dadurch gekennzeichnet, dass** ein T-Zellen Klon eingesetzt wird, um das Rezeptormolekül umfassende Material herzustellen.

5. Verfahren nach einem der vorstehenden Ansprüche, **dadurch gekennzeichnet, dass** die Membranfraktion aus Schritt 1 durch mechanische Behandlung von ganzen Zellen erhalten wird.

6. Verfahren nach einem der vorstehenden Ansprüche, **dadurch gekennzeichnet, dass** das Zelloberflächenantigen umfassende Material in das Säugetier in Abwesenheit eines Hilfsstoffs injiziert wird.

7. Verfahren nach einem der vorstehenden Ansprüche, **dadurch gekennzeichnet, dass** paramagnetische Kugeln als Träger benutzt werden.

8. Verfahren nach einem der vorstehenden Ansprüche, **dadurch gekennzeichnet, dass** die Auswahl in von mindestens einem der Schritte 6 oder 7 durchgeführt wird unter Einsatz eines Agglutination-Assays, wobei der Supernatant des B-Zellen Klon kontaktiert wird mit einem Träger, der mit Antikörpern beschichtet ist, die fähig sind, Antikörper der Art des injizierten Säugetiers zu binden, welches in Schritt 1 benutzt worden ist, und ganze Zellen, die das Zelloberflächenantigen enthalten, und Agglutination festgestellt wird.

9. Verfahren nach Anspruch 8, **dadurch gekennzeichnet, dass** ganze Zellen, die sich auf die besagten ganzen Zellen beziehen, denen es aber an dem besagten Zelloberflächenantigen ermangelt, als Kontrolle eingesetzt werden.

10. Verfahren nach einem der vorstehenden Ansprüche, **dadurch gekennzeichnet, dass** der ausgewählte B-Zellen Klon in Schritt 6 mit Myeloma-Zellen gemischt und einer Mini-Elektrofusion unterworfen wird.

11. Monoklonaler Antikörper, der mit der klonotypischen Struktur eines T-Zellen Rezeptors reagiert, **dadurch gekennzeichnet, dass** der besagte monoklonale Antikörper mit dem T-Zellen Rezeptor eines HC gp-39 reaktiven T-Zellen Klon reagiert.
- 5 12. Monoklonaler Antikörper nach Anspruch 11, **dadurch gekennzeichnet, dass** der T-Zellen Klon ECACC Zugangsnummer 96103122 ist.
13. Monoklonaler Antikörper nach Anspruch 11, **dadurch gekennzeichnet, dass** er durch ein Hybridom erzeugt wird, welches aus der Gruppe ausgewählt ist, die aus ECACC Zugangsnummer 96103118, ECACC Zugangsnummer 96103119, ECACC Zugangsnummer 96103120 und ECACC Zugangsnummer 96103121 besteht.
- 10 14. Pharmazeutische Komposition mit einem monoklonalen Antikörper nach einem der Ansprüche 11 bis 13, zugemischt mit einem geeigneten weiteren Bestandteil, geeignet für die Behandlung rheumatischer Arthritis.
- 15 15. Diagnosereagenz mit einem monoklonalen Antikörper gemäß einem der Ansprüche 11 bis 13.

Revendications

- 20 1. Procédé de préparation d'un anticorps monoclonal contre une partie déterminante de spécificité d'un antigène de surface cellulaire ayant une section constante et une section variable, dans lequel au moins une partie de ladite section variable définit une partie déterminante de spécificité dudit antigène de surface, comprenant les étapes :
- 25 1) d'injection à un mammifère non humain de matériel comprenant un antigène de surface cellulaire, ledit matériel étant choisi dans le groupe consistant en : i) cellules entières et ii) une fraction de membrane obtenue par traitement de cellules entières ;
- 30 2) d'isolement d'une fraction cellulaire contenant des lymphocytes B de la rate dudit mammifère ;
- 3) d'enrichissement de ladite fraction cellulaire obtenue à l'étape 2) en lymphocytes B spécifiques pour ledit antigène de surface cellulaire par mise en contact de la fraction cellulaire avec du matériel de cellules lié à un support correspondant auxdites cellules entières, ledit matériel de cellules correspondant lié à un support étant dépourvu dudit antigène de surface cellulaire, et séparation des lymphocytes B liés au matériel des cellules correspondantes lié à un support de la fraction cellulaire enrichie contenant des lymphocytes B non liés à utiliser à l'étape suivante ;
- 35 4) de mise en contact de la fraction cellulaire avec du matériel comprenant l'antigène de surface cellulaire lié à un support choisi dans le groupe j) de cellules entières et jj) d'une fraction membranaire obtenue à partir desdites cellules entières, et ensuite séparation des lymphocytes B non liés audit matériel lié à un support des lymphocytes B liés audit matériel lié à un support ;
- 40 5) de soumission des lymphocytes B liés audit matériel lié à un support obtenu à l'étape précédente à une dilution limitante suivie par une expansion clonale ;
- 6) de sélection d'un clone de lymphocyte B et immortalisation dudit clone de lymphocyte B sélectionné en utilisant une technique de fusion à petite échelle ; et
- 7) de sélection et clonage d'un hybridome capable de produire des anticorps qui se fixent spécifiquement audit antigène de surface cellulaire, suivis de l'isolement d'une fraction comprenant un anticorps monoclonal du surnageant dudit hybridome.
- 45 2. Procédé selon la revendication 1, **caractérisé en ce que** le mammifère ayant subi l'injection de l'antigène de surface est d'une espèce différente de l'espèce de mammifère dont est issu l'antigène de surface.
3. Procédé selon la revendication 2, **caractérisé en ce que** du matériel comprenant une molécule réceptrice est utilisé en tant que matériel comprenant l'antigène de surface.
- 50 4. Procédé selon la revendication 3, **caractérisé en ce qu'on** utilise un clone de lymphocyte T pour préparer le matériel comprenant la molécule réceptrice.
- 55 5. Procédé selon l'une quelconque des revendications précédentes, **caractérisé en ce que** la fraction membranaire à l'étape 1 s'obtient par traitement mécanique de cellules entières.
6. Procédé selon l'une quelconque des revendications précédentes, **caractérisé en ce que** le matériel comprenant

l'antigène de surface cellulaire est injecté au mammifère en l'absence d'adjuvant.

7. Procédé selon l'une quelconque des revendications précédentes, **caractérisé en ce qu'on** utilise des billes paramagnétiques en tant que support.
8. Procédé selon l'une quelconque des revendications précédentes, **caractérisé en ce que** la sélection à l'une au moins des étapes 6 et 7 est effectuée en utilisant une épreuve d'agglutination dans laquelle on met en contact le surnageant du clone de lymphocyte B avec un support recouvert d'anticorps capables de fixer des anticorps de l'espèce de mammifère ayant reçu l'injection utilisée à l'étape 1 et des cellules entières portant l'antigène de surface cellulaire, et l'agglutination est détectée.
9. Procédé selon la revendication 8, **caractérisé en ce que** des cellules entières correspondant auxdites cellules entières mais dépourvues dudit antigène de surface cellulaire sont utilisées en tant que témoin.
10. Procédé selon l'une quelconque des revendications précédentes, **caractérisé en ce que** le clone de lymphocyte B sélectionné à l'étape 6 est mélangé avec des cellules de myélome et soumis à une mini-électrofusion.
11. Anticorps monoclonal réactif avec la structure clonotypique d'un récepteur de lymphocyte T, **caractérisé en ce que** ledit anticorps monoclonal est réactif avec le récepteur de lymphocyte T d'un clone de lymphocyte T réactif avec HC gp-39.
12. Anticorps monoclonal selon la revendication 11, **caractérisé en ce que** le clone de lymphocyte T répond au numéro d'accès ECACC 96103122.
13. Anticorps monoclonal selon la revendication 11, **caractérisé en ce qu'il** est produit par un hybridome choisi dans le groupe consistant en le numéro d'accès ECACC 96103118, le numéro d'accès ECACC 96103119, le numéro d'accès ECACC 96103120 et le numéro d'accès ECACC 96103121.
14. Composition pharmaceutique comprenant un anticorps monoclonal selon l'une quelconque des revendications 11 à 13, en mélange avec un excipient approprié, appropriée au traitement de l'arthrite rhumatoïde.
15. Réactif de diagnostic comprenant un anticorps monoclonal selon l'une quelconque des revendications 11 à 13.

Figure 1: Identification of H.243 T-cell surface molecules immunoprecipitated by MAb directed against H.243.

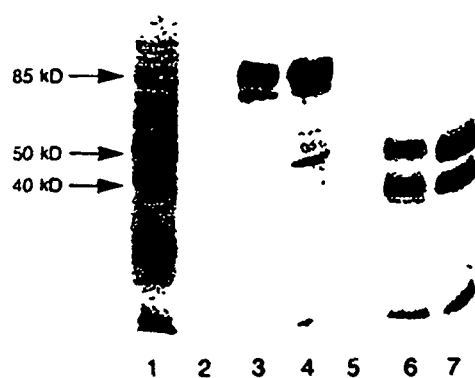
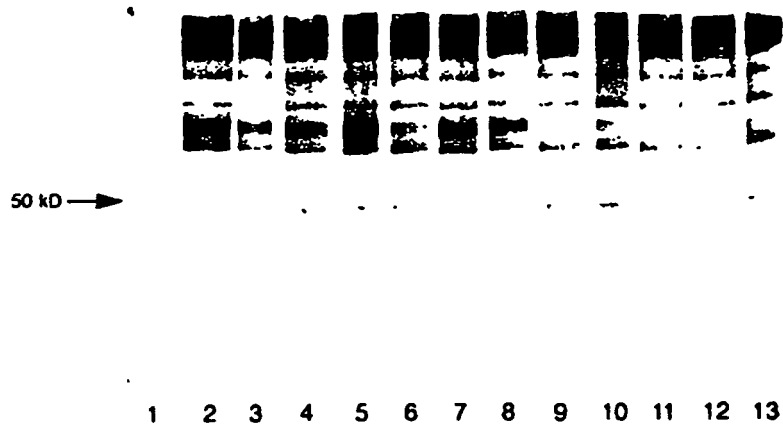


Figure 2: Anti-clonotype MAb recognize TCR $\alpha\beta$ in Western blotting.



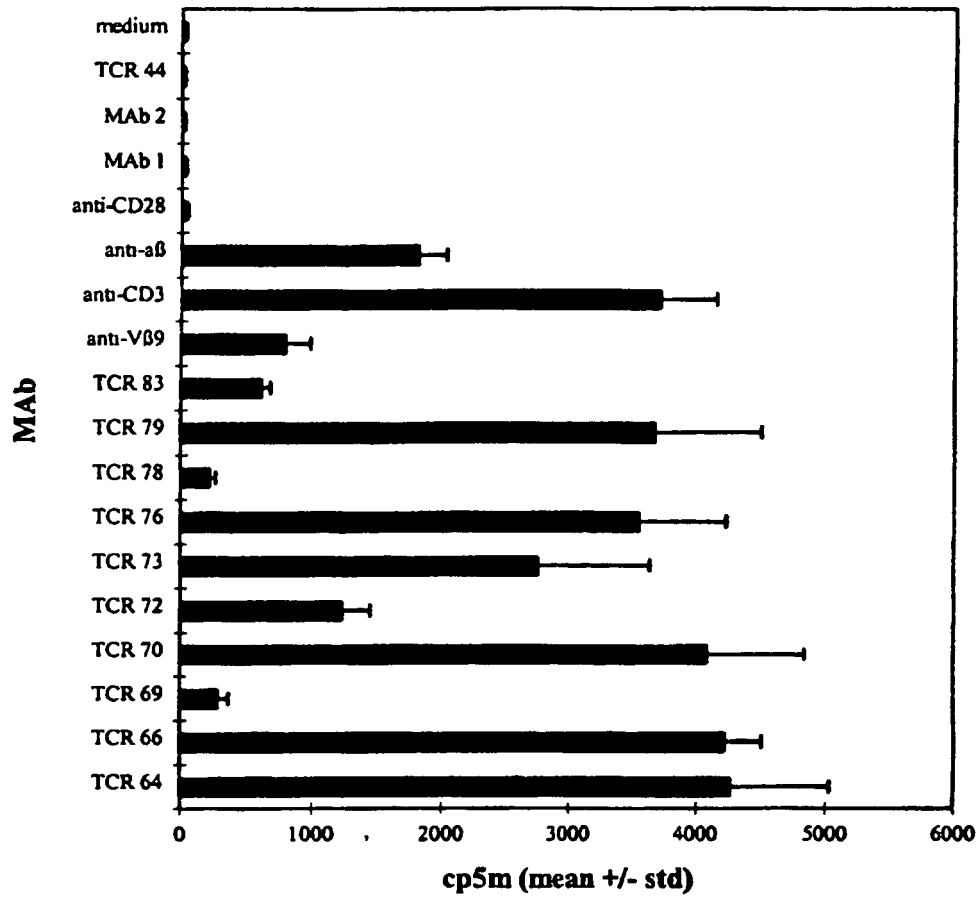
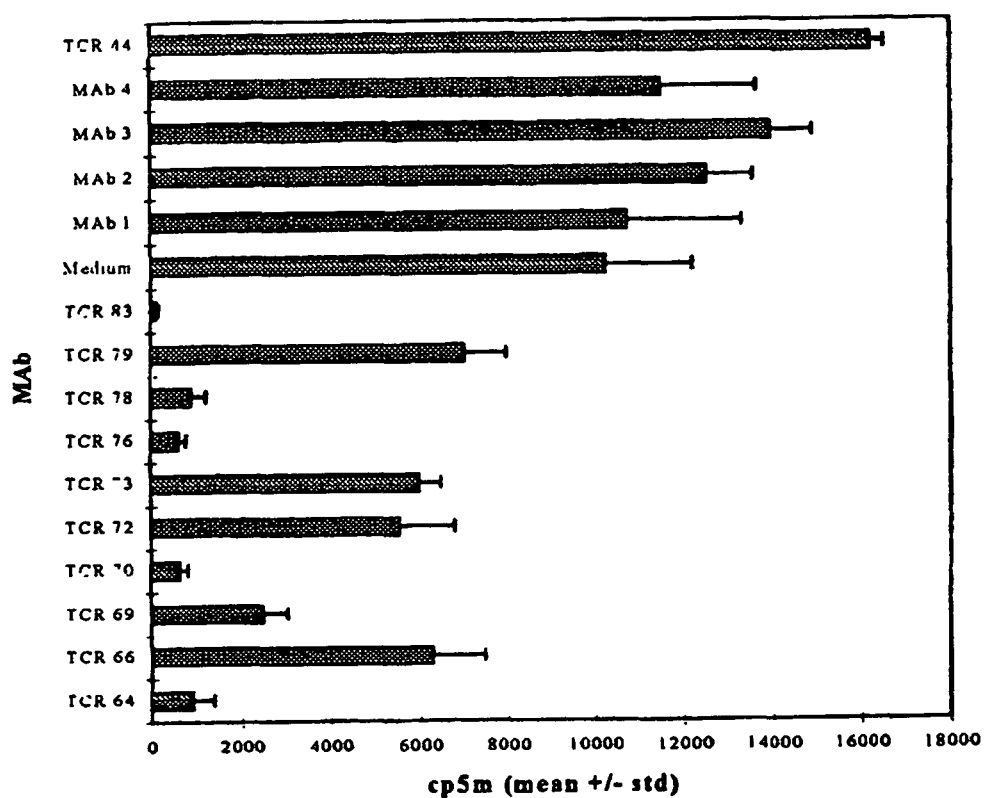
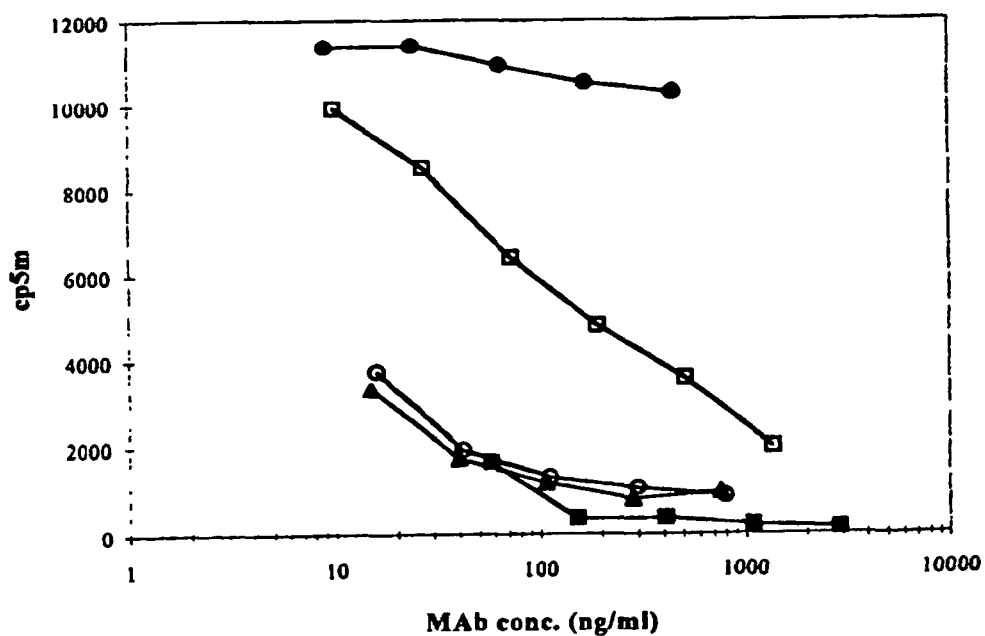


Figure 3: Immobilized anti-TCR MAb stimulate proliferation of human T-cell clone H.243.



A



B

Figure 4: Pre-incubation with anti-TCR MAb inhibit or block antigen-driven proliferation of human T-cell clone H.243.

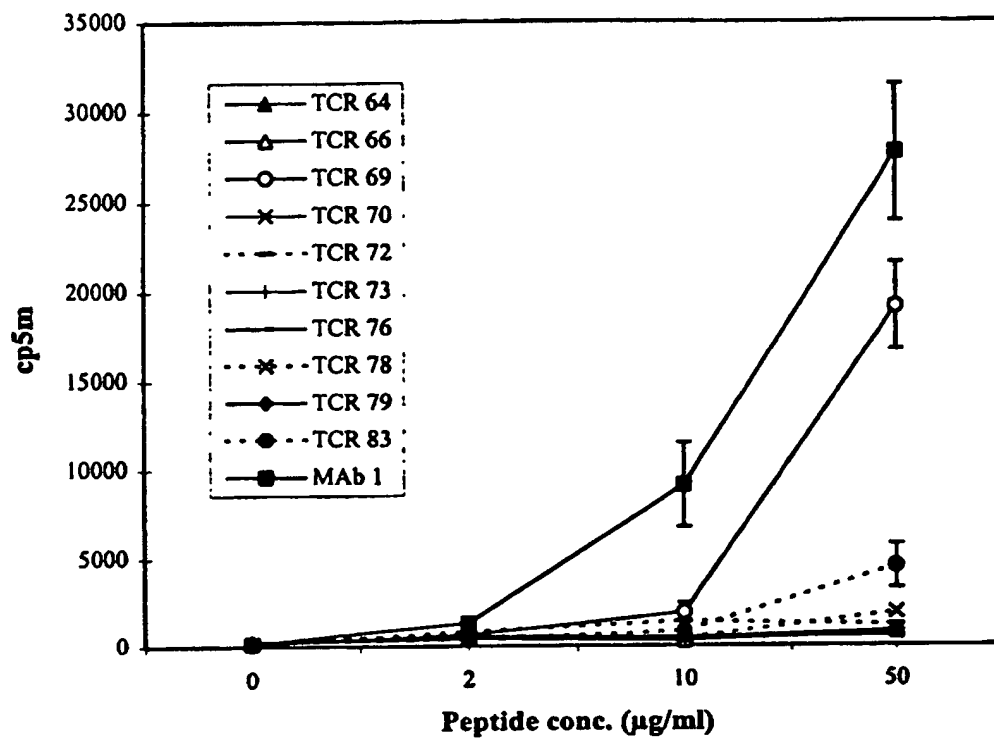


Figure 5: Anti-clonotype MAb induce energy in human Th1-clone H.243.

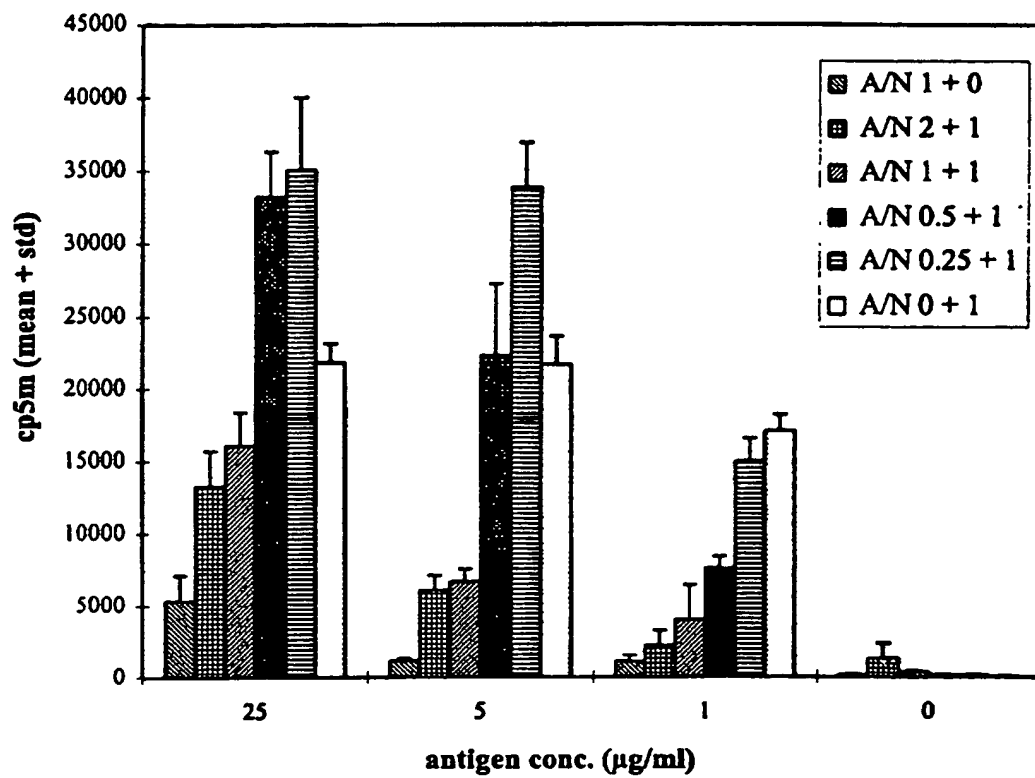


Figure 6: Anergic H.243 T cells suppress the response of non-energic H.243 T cells.